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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/644,594	08/19/2003	Tony N. Frudakis	DNA1170-2	6207
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4365 EXECU		WHALEY, PABLO S		
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

# Office Action Summary

Application No.	Applicant(s)		
10/644,594	FRUDAKIS ET AL.		
Examiner	Art Unit		
PABLO WHALEY	1631		

Office Action Gammary	Examiner	Art Unit					
	PABLO WHALEY	1631					
The MAILING DATE of this communication appears on the cover sheet with the correspondence address							
Period for Reply							
A SHORTENED STATUTORY PERIOD FOR REPLY WHICHEVER IS LONGER, FROM THE MAILING D.  - Estensions of time may be available under the provisions of 37 CFR 1.15 and 65 CFR 1.15 and	ATE OF THIS COMMUNICATION  16(a). In no event, however, may a reply be tin  till apply and will expire SIX (6) MONTHS from  cause the application to become ABANDONE	N. nely filed the mailing date of this o D (35 U.S.C. § 133).					
Status							
1) Responsive to communication(s) filed on 15 Se	eptember 2009.						
2a) This action is FINAL. 2b) ☑ This	action is non-final.						
3)☐ Since this application is in condition for allowance except for formal matters, prosecution as to the merits is							
closed in accordance with the practice under E	x parte Quayle, 1935 C.D. 11, 45	53 O.G. 213.					
Disposition of Claims							
4) Claim(s) 1 and 84-115 is/are pending in the ap	olication						
4a) Of the above claim(s) is/are periong in the application.  4a) Of the above claim(s) is/are withdrawn from consideration.							
4a) Or the above claim(s) is/are withdrawn from consideration.  5) Claim(s) is/are allowed.							
6)⊠ Claim(s) <u>1 and 84-115</u> is/are rejected.							
7) Claim(s) is/are objected to.							
8) Claim(s) are subject to restriction and/or	election requirement						
Application Papers							
9)☐ The specification is objected to by the Examine							
10)☐ The drawing(s) filed on is/are: a)☐ acce	epted or b) objected to by the I	Examiner.					
Applicant may not request that any objection to the	drawing(s) be held in abeyance. See	37 CFR 1.85(a).					
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).							
11)☐ The oath or declaration is objected to by the Ex	aminer. Note the attached Office	Action or form P	ГО-152.				
Priority under 35 U.S.C. § 119							
12) Acknowledgment is made of a claim for foreign	priority under 35 U.S.C. § 119(a)	-(d) or (f).					
a) ☐ All b) ☐ Some * c) ☐ None of:							
Certified copies of the priority documents have been received.							
2. Certified copies of the priority documents have been received in Application No							
3. Copies of the certified copies of the priority documents have been received in this National Stage							
application from the International Bureau							
* See the attached detailed Office action for a list	of the certified copies not receive	d.					
Attachment(s)							
Notice of References Cited (PTO-892)	4) Interview Summary	(PTO-413)					
2) Notice of Draftsperson's Patent Drawing Review (PTO-948)	Paper No(s)/Mail Da						
Information Disclosure Statement(s) (PTO/SB/08)  Paper No/s / Mail Date	5) Notice of Informal F 6) Other:	atert Application					

## Request For Continued Examination

A request for continued examination under 37 CFR 1.114 was filed in this application after appeal to the Board of Patent Appeals and Interferences, but prior to a decision on the appeal. Since this application is eligible for continued examination under 37 CFR 1.114 and the fee set forth in 37 CFR 1.17(e) has been timely paid, the appeal has been withdrawn pursuant to 37 CFR 1.114 and prosecution in this application has been reopened pursuant to 37 CFR 1.114. Applicant's submission filed on 09/15/2009 has been entered.

#### Status of the Claims

Claims 1 and 84-115 are pending and under consideration. Claims 2-83 are cancelled.

#### Withdrawn Rejections

The rejection of claims 1 and 83-115 under 35 U.S.C. 112, first paragraph, is withdrawn in view of applicant's amendment filed 09/15/2009.

The rejection of claims 1 and 83-115 under 35 U.S.C. 112, second paragraph, is withdrawn in view of applicant's amendment filed 09/15/2009.

The rejection of claims 1 and 83-115 under 35 U.S.C. 103(a) as being unpatentable over Parra, Shriver, Kaessmann, Vines, and Pritchard is withdrawn in view of applicant's amendment filed 09/15/2009.

### Claim rejections - 35 USC § 112, 1st Paragraph

Claims 1 and 84-115 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the written description requirement. The claims contains subject matter which was not described in the

specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor, at the time the application was filed, had possession of the claimed invention.

## This is a NEW MATTER rejection.

Claim 1 (step c) now requires that the second population of SNPs "is an autosomal marker, and wherein the at least one SNP of the second population of SNPs is not located within a region of a gene encoding a protein." While the specification does disclose autosomal markers in a Native American population [0112] and a list of autosomal chromosomes [0235], this does not provide support for the negative limitation in the claimed method step as amended. Applicant's arguments do not point to any support for this new limitation. As the newly recited limitations are not supported by the originally filed claims or disclosure, the claims are rejected for reciting new matter. This rejection is necessitated by amendment

#### Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

The factual inquiries set forth in Graham v. John Deere Co., 383 U.S. 1, 148 USPQ 459 (1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103(a) are

summarized as follows:

Determining the scope and contents of the prior art.

Ascertaining the differences between the prior art and the claims at issue.

Resolving the level of ordinary skill in the pertinent art.

4. Considering objective evidence present in the application indicating obviousness or

nonobviousness.

Claims 1, 84-86, 90-97, 99-100, and 104-109 are rejected under 35 U.S.C. 103(a) as being unpatentable over Parra et al. (Am. J. Physical Antropol., January 2001, Vol. 114, Issue 1, p. 18-29), in view of Cargill et al. (Nature Genetics, 1999, Vol. 22, p.231-238).

The amended claims are now drawn to a method of inferring, with a predetermined level of confidence, proportional ancestry of at least two ancestral groups of a test individual by identification of a population structure comprising: a) determining single nucleotide polymorphisms (SNPs) for a first population and identifying a first population of SNPs having a frequency differential (delta) > 0.4 between one or more pairs of population groups wherein the first population of SNPs are identified from a database in silico; b) contacting a parental sample nucleic acid with one or more hybridizing nucleic acids corresponding the first population of SNPs, wherein the one or more hybridizing nucleic acids selectively hybridize to the nucleic acid in the parental sample; c) selecting SNPs hybridizing in step (b) to generate a second population of SNPs which have a minor allele frequency > 1% and (delta) > 0.4 for at least one pair of the at least two population groups, wherein at least one of the second population of SNPs is a SNP which may be correlated with but not linked to a gene-linked trait, wherein the second population of SNPs is an autosomal marker, and wherein the at least one SNP of the second population of SNPs is not located within a region of a gene encoding a protein; d) contacting a sample comprising nucleic acid molecules of a test individual with at least 20 second population of SNPs, wherein the second population of SNPs are indicative of a population structure, and wherein the population structure is correlated with a trait of the test individual; e) determining the nucleotide occurrences of the second population of SNPs in the sample

from the test individual; and f) identifying the population structure indicated by the nucleotide occurrences determined for the test individual, wherein identifying the population structure infers the proportional ancestry of the test individual; and g) providing information resulting from steps (a) through (f) to a user.

Parra teaches a method for inferring the extent of European admixture in six different African-American populations [See Abstract]. In particular, Parra teaches a method for identifying a population structure by determining SNP markers for a first population using a battery of autosomal markers (both linked and unlinked) [See Abstract, and p.19-20, DNA Analysis Section, and Table 1]. Parra identifies allele frequencies corresponding to the 10 autosomal markers have frequencies differentials > 0.4 between one or more populations [Table 1, last column], which meets the claim language. The method requires standard PCR genotyping procedure, which inherently requires contacting parent samples with markers [See p.20, Col. 1 and Table 1, last column]. In this population of markers, at least one is an autosomal polymorphism marker (D11S429) [p.20, Col. 1, and Table 1] that is associated with ancestry [Table 5], which shows the use at least one autosomal SNP that may be correlated but not linked to a gene-trait. A combination of SNP markers are selected to obtain an estimate of admixture for a subpopulation [p.21, Col. 1, ¶ 2, and Fig. 1], wherein allele frequencies of the SNP markers are > 1% [Table 1], which is a teaching for minor allele frequencies. Parra shows the use of markers that are unlinked to certain loci [p.20, Col. 1, and p.23, Col. 2]. Parra calculates the frequency differences between populations based on SNPs [p.23, Col. 2]. The admixture distribution of the non-parental test populations is inferred within a predetermined confidence interval [p. 21 and Table 2]. Parra also shows fitting genotype frequencies to Hardy-Weinberg proportions and suggest the selection of genetic markers that show homogeneity with Africa and Europe based on allele frequency [p.20, Statistical Analysis]. Parra discloses a biogeographical ancestry trait (BGA) [Fig. 1], and admixture proportions of samples estimated using maximum likelihood calculations [p.20, Col. 2, ¶2 and ¶3]. Parra shows proportional ancestry comprising a three-way comparison of sub-populations of African-Americans and the distribution percentage of European alleles within this sub-population [Fig. 1] derived from maximum-likelihood methods [p.25, Col. 2]. Parra shows SNPs detected in a subpopulation of non-parental individuals for determining sub-population structure [Fig. 1].

Parra does not teach a first population of SNPs identified from a database in silico, as in claim 1.

Parra does not teach selecting SNPs from a first population to generate a second population of SNPs wherein the at least one SNP of the second population of SNPs have minor allele frequencies >1% and are not located within a region of a gene encoding a protein, as in claim 1.

Cargill teaches a method for screening samples to determine whether identified SNPs are coding or non-coding SNPs using well known genotyping techniques [p.232-233, Results, Table 1, and Table 2]. The method includes collecting gene sequence data from known public databases and suggests construction of comprehensive SNP databases to improve screening [See p. 237, Methods and p.236, Col. 1st paragraph]. Cargill also determines the distribution of minor allele frequency, wherein SNPs are classified according to whether minor allele frequency is high (>15%), medium (5-15%), or low (<1%) [p.234, Col. 1, last para., and Col. 2, and Fig. 1], which meets the claim limitation of minor allele frequencies > 1%.

It would have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method of Parra by selecting a first population of SNPs identified from a database in silico, as in claim 1, since Parra suggests genetic markers are easily obtained [p.20, Col. 1] and since Cargill shows computations methods for collecting gene sequence data from known public databases and suggests construction of comprehensive SNP databases [See p. 237, Methods and p.236, Col. 1<sup>st</sup> paragraph]. The motivation would have been to improve screening—genotyping analysis by using comprehensive databases containing data from many ethnic groups, as suggested by Cargill [p.236, Col. 1<sup>st</sup> paragraph].

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It would have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method of Parra by screening the first population of SNPs to generate a second population of SNPs wherein the at least one SNP of the second population of SNPs is not located within a region of a gene encoding a protein, as in claim 1, since Cargill shows screening genetic samples to determine whether identified SNPs are coding or non-coding SNPs that occur outside of the coding regions of a gene [p.232-233, Results, Table 1, and Table 2]. The motivation would have been to obtain the most informative SNP markers by screening the set of identified SNPs for non-coding SNPs which are known to effect biological function [Cargill, p.236, Col. 1, last paragraph].

Claims 1, 84-97, 99-100, and 104-115 are rejected under 35 U.S.C. 103(a) as being unpatentable over Parra et al. (Am. J. Physical Antropol., January 2001, Vol. 114, Issue 1, p. 18-29), in view of Cargill et al. (Nature Genetics, 1999, Vol. 22, p.231-238), and further in view of Sorenson et al. (US 2003/0172065; Filed Mar. 29, 2002).

Parra and Cargill make obvious the method of claims 1, 84-86, 90-97, 99-100, and 104-109, as set forth above.

Parra and Cargill do not specifically teach contacting samples with high numbers of SNPs, as in claims 87-89.

Parra and Cargill do not teach proportional ancestries comprising a photo of a person from whom the known proportional ancestry was determined, as in claims 110-115.

Sorenson discloses a genealogical research and record keeping system for identifying commonalities in haplotypes from biological samples [Abstract]. In particular, Sorenson teaches thousands of known genetic markers and millions of characterized SNPs may be analyzed [0042], [Fig. 4] for identifying a population structure [0032, 0046-0047], as in claim 87-89. Sorenson also discloses prior

art genetic records of human eye, hair and skin color, height and other physical characteristics [0009], and ancestral data stored on microfiche and on a number of other electronic media formats including the internet [0003], which is broadly interpreted as a teaching for digital information and pictures as in claims 110-115.

It would have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method made obvious by Parra and Cargill by additionally contacting samples with high numbers of SNPs as in claims 87-89, since Sorenson teaches genetic analysis using thousands of known genetic markers and millions of characterized SNPs [0042, Fig. 4]. The motivation would have been to address the need for correlating genetic information with genealogical information to identify previously unknown biological relationships or previous family ties, as suggested by as suggested by Sorenson [0013, 0015].

It would further have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method made obvious by Parra and Cargill by using proportional ancestries comprising a photo of a person from whom the known proportional ancestry was determined, as in claims 110-115, since Sorenson teaches genetic records and ancestral data stored on microfiche and in electronic media format [0003, 0009]. The motivation would have been to improve the sharing of ancestral data using electronic formats that are suitable for the internet, as suggested by Sorenson [0003].

Claims 1, 84-101, and 104-115 are rejected under 35 U.S.C. 103(a) as being unpatentable over Parra et al. (Am. J. Physical Antropol., January 2001, Vol. 114, Issue 1, p. 18-29), in view of Cargill et al. (Nature Genetics, 1999, Vol. 22, p.231-238), in view of Sorenson et al. (US 2003/0172065; Filed Mar. 29, 2002), and further in view of Shriver et al. (American Journal of Human Genetics, 1997, Vol. 60, p.957-964; IDS filed 5/20/2004).

Parra, Cargill, and Sorenson make obvious the method of claims 1, 84-97, 99-100, and 104-115, as set forth above. Parra also shows two-way and three-way comparison of populations that are both intracontinental and intercontinental [Table 1, Fig. 1, and p. 22].

Parra, Cargill, and Sorenson do not teach performing a likelihood determination for affiliation with an East Asian ancestral group, as in claim 98.

Parra, Cargill, and Sorenson do not teach performing four-way comparisons, as in claim 101.

Shriver teaches a method for identifying a set of genetic markers using likelihood analysis that allows the confident determination of ethnicity for use in a forensic setting [Abstract and p.964, Discussion]. In particular, Shriver presents population specific alleles (PSAs) [p.957, Col. 2], as well as methods for calculating allele-frequency differentials between test samples of different populations [p.958, Col. 2] and for calculating likelihood values for different loci [Table 1, and Table 2]. Shriver does not specifically teach "four-way" comparison. However, Shriver performs two-way and three-way comparisons between multiple populations [Fig. 1-4]. Therefore, it would be obvious to one of ordinary skill in the art to perform a four-way comparison among four ancestral groups, as in claim 101. Shriver also suggests similar markers could be developed for the identification of other populations including those of Asian origin [p.963, last ¶, Col. 1].

It would further have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method made obvious by Parra, Cargill, and Sorenson by performing a likelihood determination for affiliation with an East Asian ancestral group as required by claims 98, since Shriver identifies genetic markers using likelihood analysis [Abstract and p.964, Discussion, Table 1, and Table 2] and suggests markers for those of Asian ancestry [p.963, last ¶, Col. 1]. The motivation would have been to allow for the confident determination of ethnicity in forensic settings, as suggested by Shriver [Abstract and p.964, Discussion, Table 1, and Table 2].

It would further have been obvious to one of ordinary skill in the art at the time of the instant invention to modify the method made obvious by Parra, Cargill, and Sorenson by performing four-way comparisons, as in claim 101, since Parra and Shriver show performing two-way and three-way comparisons between multiple populations, as set forth above. The motivation would have been to identify other populations that share common ancestry, as suggested by Shriver [p.963, Col. 1, last ¶, and Col. 2].

Claims 1 and 84-115 are rejected under 35 U.S.C. 103(a) as being unpatentable over Parra et al. (Am. J. Physical Antropol., January 2001, Vol. 114, Issue 1, p. 18-29), in view of Cargill et al. (Nature Genetics, 1999, Vol. 22, p.231-238), in view of Sorenson et al. (US 2003/0172065; Filed Mar. 29, 2002), in view of Shriver et al. (American Journal of Human Genetics, 1997, Vol. 60, p.957-964; IDS filed 5/20/2004), and further in view of Pritchard et al. (Theoretical Population Biology, 2001, Vol. 60, p. 227-237).

Parra, Cargill, Sorenson, and Shriver make obvious the method of claims 1, 84-101, and 104-115, as set forth above.

Parra, Cargill, Sorenson, and Shriver do not teach generating a graphical representation of the comparison of three ancestral groups, wherein the representation comprises a confidence contour, as in claims 102 and 103.

Pritchard teaches a method for inferring proportional ancestry of different ancestral groups in a population structure using a graphical display format [Fig. 1], as in claims 102 and 103. Points in the extreme corners of the triangular plots are correctly classified [Fig. 1], which shows groups represented in a vertex of a triangle. Ancestry is also represented using a line plot, wherein dashed lines represent individuals from a population with the most divergent allele frequencies [Fig. 1 and p.232, Col. 2], which suggests a contour indicating a level of confidence. Pritchard also teaches a computer-based program

STRUCTURE for estimating population structure for 20 data sets of 50, 200, and 1000 biallelic markers [p. 232, Results].

It would have been obvious to one of ordinary skill in the art at the time of the instant invention to practice the method made obvious by Parra, Cargill, Sorenson, and Shriver by generating a graphical representation, as in claims 102 and 103, since Pritchard shows graphically displaying ancestral results in triangular format [Fig. 1]. The motivation would have been to use a user-friendly graphical method for inferring ancestry in a plurality of populations, as suggested by Pritchard [Fig. 1].

## Response to Arguments

Applicant's arguments, filed 09/15/2009, have been fully considered but are moot in view of the new grounds of rejection.

#### Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Pablo Whaley whose telephone number is (571)272-4425. The examiner can normally be reached between 12pm-8pm.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Marjorie Moran can be reached at 571-272-0720. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

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Art Unit: 1631

Pablo S. Whaley

Patent Examiner

Art Unit 1631

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Primary Examiner, Art Unit 1631